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## PATENT COOPERATION TREATY

# **PCT**

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### INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Artcle 36 and Rule 70)

Applicant's or agent's file reference 2FPO-02-09	FOR FURTHER ACTION	SeeNotification Examination I	nofTransmittalofInternationa Report (Form PCT/IPEA/416	alPreliminary
International application No. PCT/KR2002/000595	International filing date(day/med) APRIL 2002 (03.04.2		Priority date (day/month/ye	ear)
International Patent Classification (IPC) IPC7 A61K 31/47	or national classification and IP	c		
Applicant  CHEMON INC. et al				
This international preliminary e     and is transmitted to the applicant.	nt according to Article 36.			ning Authority
amended and are the basis	of sheets, included an included by ANNEXES, i.e., sheet for this report and/or sheets could be Administrative Instructions under the Instruction u	s of the description taining rectificat	on, claims and/or drawings v	which have been nority (see Rule
These annexes consist of a tota	of 10 sheets.			
IV Lack of unity of in  V X Reasoned statem citations and expl  VI Certain document  VII Certain defects in	t of opinion with regard to novel nvention ent under Article 35(2) with regard anations supporting such statements cited the international application ons on the international application	rd to novelty, invent	entive step or industrial appli	icability;
Date of submission of the demand 03 NOVEMBER 2003 (03.11)		ate of completion  27 MAY 20	of this report 004 (27.05.2004)	
Name and mailing address of the IPE Korean Intellectual Prop 920 Dunsan-dong, Seo-g Republic of Korea	erty Office gu, Daejeon 302-701,	LEE, HYUN		
Facsimile No. 82-42-472-7140	ĮΤ	elephone No. 82	-42-481-5606	



International aplication No.

PCT/KR2002/000595

#### I. Basis of the report With regard to the elements of the international application:\* the international application as originally filed the description: , as originally filed pages 2-5, 9-12, 15-39, 41-49 , filed with the demand pages , filed with the letter of 12/05/2004 pages 1, 6, 7, 8, 13, 14, 40 the claims: , as originally filed pages , as amended (together with any statment) under Article 19 pages , filed with the demand pages , filed with the letter of 12/05/2004 pages <u>50-52</u> the drawings: , as originally filed pages \_ , filed with the demand pages \_ \_\_\_\_\_ filed with the letter of pages \_ the sequence listing part of the description: , as originally filed pages \_ , filed with the demand pages , filed with the letter of pages 2. With regard to the language, all the elements marked above were available or furnished to this Authority in the language in which the international application was filed, unless otherwise indicated under this item. which is These elements were available or furnished to this Authority in the following language English the language of a translation furnished for the purposes of international search (under Rule 23.1(b)). the language of publication of the international application (under Rule 48.3(b)). the language of the translation furnished for the purposes of international preliminary examination (under Rules 55.2 and/ or 55.3). 3. With regard to any nucleotide and/or amino acid sequence disclosed in the international application, the international preliminary examination was carried out on the basis of the sequence listing: contained in the international application in written form. filed together with the international application in computer readable form. furnished subsequently to this Authority in written form. furnished subsequently to this Authority in computer readable form

The amendments have resulted in the cancellation of: the description, pages

international applicationas as filed has been furinshed.

the claims, Nos. the drawings, sheet

been furnished.

5.

This report has been established as if (some of) the amendments had not been made, since they have been considered to go beyond the disclosure as filed, as indicated in the Supplemental Box(Rule 70.2(c)).\*\*

The statement that the subsequently furnished written sequence listing does not go beyond the disc losure in the

The statement that the information recorded in computer readable form is identical to the written sequence listing has

Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this opinion as "originally filed." and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17).

\*\* Any replacement sheet containing such amendments must be referred to under item I and annexed to this report.

### INTERNATIONAL PRELIMINARY EXAMINATION

International aplication No.

PCT/KR2002/000595

V.	V. Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement			
1.	Statement			A
	Novelty (N)	Claims	1-6	YES
	21012107 (217)	Claims		NO
	Inventive step (IS)	Claims	1-6	YES
	mvenuve step (15)	Claims		NO
	Industrial applicability (IA)	Claims	1-6	YES
		Claims		NO
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2. Citations and explanations (Rule 70.7)

The following documents are referred to:

D1: US-A-4943577 D2: US-A-5372813 D3: US-A-6166205

- 1. D1 discloses a piperazinyl quinoline compound having an alkyl piperazinyl side chain for the treatment of a mental disorder. D3 discloses a piperazinyl quinoline compound having an imidazole side chain useful as a dopamine (D4) antagonist. Both D1 and D3 have the same structural moiety but have differential side chains, compared to the present claims 1-6. The most relevant compound disclosed in D2 cannot destroy the novelty of claims 1-6 because of the claims amended by the letter of 12/05/2004. Accordingly, the subject matter of claims 1-6 seems to be novel (Article 33(2) PCT).
- 2. The closest state of the art appears to be represented by D2 which discloses a method for the measurement of serotonin uptake sites in a sample using piperazinyl chemical structural difference between D2 and the present application quinoline. The is the position of the side chain on quinoline. The compound in D2 has side chains at 3, 5, 7, and 8 positions of quinoline, while the present compound has them at 3, 4, and 6 positions. The addition of the side chain to quinoline moiety does not require a special reaction scheme. Consequently, the inventive step of the present application should be considered based on the effect resulting from the substitution of side chains on quinoline. Compared to D2, the present compounds show an outstanding affinity for the serotonin transfer (e.g. the affinity of the compound shown in Example 10 is 10 times as high as that of the compound with H at 4 position of quinoline). Consequently, the inventive step of claims 1-3 can be acknowledged. The subject matter of claims 4-6 is a method for preparing a compound and a pharmaceutical usage. The inventive step of claims 4-6 can be acknowledged because the inventive step of the present compound is approved.

Thus, claims 1-6 involve an inventive step and meet the requirement of Article 33(3) PCT.

The present application is considered to meet the criteria of industrial applicability (Article 33(4) PCT).

# Rec'd POTATTO 27 SEP 2004

# QUINOLINE DERIVATIVES, THEIR PREPARATION AND PHARMACEUTICAL COMPOSITIONS COMPRISING THE SAME

### TECHNICAL FIELD

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The present invention relates to quinoline derivatives represented by in formula 1 as below. More specifically, the present invention relates to quinoline derivatives and their salts that pharmaceutically acceptable interrupt reuptake of serotonin into presynaptic neuron and thus increase the concentration of serotonin in synapse. present invention also includes the process for preparing the said compounds of formula 1 and their pharmaceutical compositions to prevent or treat serotonin-related mental effective said compounds as disorders comprising the ingredients.

Formula 1

$$R^4$$
  $R^3$   $R^4$   $R^4$ 

wherein,

 $R^1$  is piperazinyl, 2-methylpiperazinyl, perhydrodiazepinyl or N-methyl-N-(2-N'-methylamino)ethylamine;

 $R^2$  is H, halogen atom,  $C_1 \sim C_4$  alkyl or  $C_1 \sim C_4$  haloalkyl;

R3 is H, halogen atom, vinyl or furanyl group; and

 ${\ensuremath{\mathsf{R}}}^4$  is halogen atom or nitro group.

### BACKGROUND ART

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nitro-2-(piperazin-1-yl)quinoline is administered to depression-induced mice and 0.5% dimethylsulfoxide (DMSO) is administered to them as a control;

Figure 2B is a graph illustrating the anti-depression effect obtained by measuring of immobility time using a tail suspension test wherein 1 mg/kg or 10 mg/kg of 4-chloro-6-nitro-2-piperazine is administered to depression-induced mice and 0.5% dimethylsulfoxide (DMSO) is administered to them as a control.

### DISCLOSURE OF INVENTION

The present invention provides quinoline derivatives and their pharmaceutical acceptable salt represented by the following formula 1:

Formula 1

wherein,

 $R^1$  is piperazinyl, 2-methylpiperazinyl, perhydrodiazepinyl or N-methyl-N-(2-N'-methylamino)ethylamine group;

 $R^2$  is H, halogen atom,  $C_1 \sim C_4$  alkyl or  $C_1 \sim C_4$  haloalkyl;  $R^3$  is H, halogen atom, vinyl or furanyl group; and  $R^4$  is halogen atom or nitro group.

More preferably,

 $R^{1}$  is 2-methylpiperazinyl, perhydrodiacepinyl, is  $N^{2}$ 



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methyl-N-(2-N'-methylamino) ethylamine group;

 ${\ensuremath{\mbox{R}}}^2$  is H, bromine, methyl, ethyl, propyl, chloropropyl or fluoropropyl group;

 $\mathbb{R}^3$  is H, chorine, bromine, iodine, vinyl or 2-furanyl group; and

R4 is chlorine, bromine or nitro group.

Most preferably, examples of the compounds represented by the chemical formula 1 include the following table 1:

Table 1

Table 1				
Example	Compound	Formula		
1	3-methyl-6-nitro-2-(piperazin-1- yl)quinoline	O <sub>2</sub> N CH <sub>3</sub>		
2	3-ethyl-6-nitro-2-(piperazin-1- yl)quinoline	O <sub>2</sub> N CH <sub>3</sub>		
3	6-nıtro-2-(piperazin-1-yl)-3- propylquinoline	O <sup>3</sup> N CH <sup>3</sup>		
4	3-(3-chloropropyl)-6-nitro- (piperazin-1-yl)quinoline	O <sub>2</sub> N CI		
5	6-iodo-2-(piperazin-1-yl)quinoline	I N N N N N N N N N N N N N N N N N N N		
6	6-bromo-2-(piperazin-1- yl)quinoline	Br. NH		
7	6-chloro-2-(piperazin-1- yl)quinoline	CI		

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8	3-(3-fluoropropyl)-6-nitro-2- (piperazin-1-yl)quinoline	O <sub>2</sub> N F
9	3-bromo-6-nitro-2-(piperazin-1- yl)quinoline	N N N N N N N N N N N N N N N N N N N
10	4-chloro-6-nitro-2-(piperazin-1- yl)quinoline	O <sub>2</sub> N CI
11	4-bromo-6-nitro-2-(piperazin-1- yl)quinoline	O <sub>2</sub> N N N N N N N N N N N N N N N N N N N
12	4-iodo-6-nitro-2-(piperazin-1- yl)quinoline	O <sub>2</sub> N N N N N N N N N N N N N N N N N N N
13	6-nitro-2-(piperazin-1-yl)-4- vinylquinoline	02N
14	4-(2-furanyl)-6-nitro-2- (piperazin-1-yl)quinoline	O <sub>2</sub> N , , , , , , , , , , , , , , , , , , ,
15	2-(3-methylpiperazin-1-yl)-6- nitroquinoline	02N
16	2-(N-methyl-N-(2-N'- methylamino)ethyl)amino-6- nitroquinoline	O <sub>2</sub> N V V V V V
17	2-perhydrodiazepin-1-yl-6- nitroquinoline	O <sub>2</sub> N

The present invention further includes scluated compounds and hydrates prepared by use of the quincline derivatives of formula 1 and their pharmaceutically acceptable salts.



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tributyl(vinyl)tin or tributyl(2-furanyl)tin in the presence of palladium catalyst to introduce vinyl or furanyl at 4-position of compounds of formula 6; and

obtained mixture with acid b) treating thus compounds.

Preferably, the Stille reaction is carried out from 90 °C to 120 °C under inert gas, e.g.,  $N_2(q)$ .

In this case, it is possible to prepare the compounds substituted with aryl, heteroaryl and aryl group besides vinyl and furanyl group.

In accordance with yet another aspect of the present invention, there is provided a method for preparing an 2-(3-methylamino) ethylamino-6-nitroquinoline and 2perhydrodiazepin-1-yl-6-nitroquinoline, represented by the following reaction scheme 4:

Reaction Scheme 4

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$$O_2N$$
 $R^1H$ 
 $O_2N$ 
 $R^1$ 
 $R$ 

wherein,

2-methylpiperazinyl, N-methyl-N-(2-N'-R1 is methylamino; ethylamine or perhydrodiazecinyl group.



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As shown, the compounds of formula 9 including an 2-(3-methylpiperazin-1-yl)-6-nitroquinoline, 2-(N-methyl-N-(2-N'-methylamino)ethylamino-6-nitroquinoline and 2-perhydrodiazepin-1-yl-6-nitroquinoline is prepared by reacting 2-chloro-6-nitroquinoline of formula 8 with 2-methylpiperazine, N-methyl-N-(2-N'-methylamino)ethylamine or perhydrodiazepine.

In accordance with a further aspect of the present invention, there is provided a pharmaceutical composition comprising the compounds of formula 1 as an effective ingredient to prevent or treat mental disorder caused by serotonin.

Concretely, the quinoline derivatives of the invention may be utilized to prevent or treat mental disorders, especially to depression.

In a preferable embodiment of the present invention, the brain tissue was gently isolated from a mouse, grounded, and then the biological activity against SERT was measured in a variety of concentration. As a result, the compounds of the present invention shows a much higher Ki value than that of the already commercialized Flucketine and a similar value to that of Paroxetine. As shown table 3, especially, 4-chloro-6-nitro-2-piperazin-1-yl-quinoline shows an excellent binding affinity over ten times than 6-nitro-2-

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8.0 Hz, 1H), 4.08 (t, J = 6.9 Hz, 2H), 3.24-3.38 (m, 5H), 2.68 (s, 3H).

 $^{13}\text{C}$  NMR (D<sub>2</sub>O)  $\delta$  153.2, 143.5, 143.2, 139.5, 126.2, 124.3, 119.9, 118.6, 112.6, 47.9, 44.6, 38.1, 32.7.

# EXAMPLE 17 : Preparation of 2-perhydrodiazepin-1-yl-6-nitroquinoline

In 10 ml of DMF, 2-chloro-6-nitroquinoline (500 mg, 2.40 mmol) was dissolved, added perhydrodiazepine (894 mg, 7.20 mmol) at room temperature, and then reacted at 80 °C for 5 hours. After the completion of the reaction, the mixture was cooled to room temperature, added water to produce a precipitate. The precipitate was isolated by filtration, washed with 50 ml of water, and then dried over for 3 hours in vacuo to give the desired product as a yellowish solid (602 mg, 2.21 mmol, 92%).

 $^{1}$ H NMR (CDCl<sub>3</sub>)  $\delta$  8.52 (d, J = 2.6 Hz, 1H), 8.27 (dd, J = 9.3, 2.7 Hz, 1H), 7.93 (d, J = 9.2 Hz, 1H), 7.62 (d, 9.4 Hz, 1H), 6.97 (d, J = 9.0 Hz, 1H), 3.87-3.93 (m, 4H), 3.11 (t, J = 5.3 Hz, 2H), 2.89 (t, J = 5.9 Hz, 2H), 2.22 (br s, 1H), 1.91-2.03 (m, 2H);

<sup>13</sup>C NMR (DMSO-d<sub>6</sub>) δ 157.1, 150.5, 139.4, 137.7, 125.4, 25 123.6, 122.2, 119.7, 46.92, 46.03, 45.41, 37.2, 27.5.



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## WHAT IS CLAIMED IS

1. A quinoline derivative of formula 1, or pharmaceutically acceptable salt of the same:

formula 1

$$R^4$$
  $R^3$   $R^2$ 

wherein,

 $R^1$  is piperazinyl, 2-methylpiperazinyl, perhydrodiazepinyl or N-methyl-N- 2-N'- methylamino)ethylamine group;

 $R^2$  is H, halogen atom,  $C_1 \sim C_4$  alkyl or  $C_1 \sim C_4$  halcalkyl;

R<sup>3</sup> is H, halogen atom, vinyl or furanyl group; and

R4 is halogen atom or nitro group

with the proviso that

when R1 is piperazinyl or 2-methylpiperazinyl;

R<sup>3</sup> is H; and

R4 is nitro group,

 $R^2$  is not H, halogen atom,  $C_1{\sim}C_3$  alkyl, or  $C_1{\sim}C_2$  haloalkyl.

2. The derivative of claim 1, wherein  $R^2$  is 2-methylpiperazinyl, perhydrodiazepinyl or N-methyl-N-2-N'-methylamine;

 $R^2$  is H, bromine, methyl, ethyl, thloropropyl or fluoropropyl:

 ${\ensuremath{R^3}}$  is H, chlorine, bromine, iodine, vinyl or 2-furanyl group; and

 $R^4$  is chlorine, bromine, iodine, or nitro group with the proviso that when  $R^1$  is 2-methylpiperazinyl;  $R^3$  is H; and  $R^4$  is nitro group,  $R^2$  is not H, bromine, methyl or ethyl.

- 3. The derivative of claim 1, wherein the derivative is selected from the group consisting of:
  - 3-(3-chloropropyl)-6-nitro-2-piperazin-1-yl-quinoline;
  - 3-(3-fluoropropyl)-6-nitro-2-piperazin-1-yl-quinoline;
  - 6-iodo-2-pipeerazin-1-yl-quinoline;
  - 6-bromo-2-piperazine-1-yl-quinoline;
- 10 6-chloro-2-piperazin-1-yl-quinoline;

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- 4-chloro-6-nitro-2-piperazin-1-yl-quinoline;
- 4-bromo-6-nitro-2-piperazin-1-yl-quinoline;
- 4-iodo-6-nitro-2-piperazin-1-yl-quinoline;
- 6-nitro-2-piperazin-1-yl-4-vinylquinoline;
- 4-(2-furanyl)-6-nitro-2-piperazin-1-yl-quinoline;
  - 2-(N-methyl-N-(2-N'-methylamino)ethyl)amino-6-
  - nitroquinoline; and
  - 2-perhydrodiazepin-1-yl-6-nitoquinoline.
- 20 4. A method for preparing a compound of formula 3, which



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comprises substituting a quinoline compound of formula 2 with piperazine to introduce a piperazinyl group at 2-position of the quinoline compound of formula 2:

Reaction Scheme 1

wherein,

 $R^2$  is H, halogen atom,  $C_1 \sim C_4$  alkyl or  $C_1 \sim C_4$  haloalkyl;  $R^3$  is H, halogen atom, vinyl or furanyl group; and  $R^4$  is halogen atom or nitro group.

- 5. A pharmaceutical composition comprising the quinoline derivative of the formula 1 as an effective ingredient for preventing or treating serotonin-related mental disorder.
- 15 6. The composition of claim 1, wherein the mental disorder is a depression.